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(54) BINDER RESIN COMPOSITIONS, PRODUCTION PROCESS, AND USE THEREOF

(57) The invention provides a novel binder resin composition with good adherence onto prime materials of polyolefin, poly(vinyl chloride), polycarbonate, PET, ABS and nylon, and also with excellent solvent solubility.

A binder resin composition characterized by being chlorinated propylenic random copolymer with weight average molecular weight of 3000 to 250000, wherein propylenic random copolymer with melting point (Tm) measured by differential scanning calorimeter (DSC) of 115 to 165°C obtained by copolymerizing propylene with other α -olefin in the coexistence of metallocene type catalyst is chlorinated to chlorine content of 10 to 40% by weight, after thermal degradation or directly without thermal degradation, and/or a binder resin com-

position characterized by containing carboxyl group-containing chlorinated propylenic random copolymer with weight average molecular weight of 30000 to 220000, wherein propylenic random copolymer with melting point (Tm) measured by differential scanning calorimeter (DSC) of 115 to 165°C obtained by copolymerizing propylene with other α -olefin in the coexistence of metallocene type catalyst is grafted with α,β -unsaturated carboxylic acid or its anhydride in amounts of 0.1 to 20% by weight, after thermal degradation or directly without thermal degradation, and then chlorinated to chlorine content of 10 to 40% by weight, stabilizer and organic solvent.

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Description

Technical field

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[0001] The present invention relates to a binder resin composition to be used for the purpose of protection or beautiful ornament of polyolefinic resins such as polypropylene, polyethylene, ethylene-propylene copolymer and ethylene-propylene-diene copolymer, vinyl chloride resin, polycarbonate (PC) resin, poly(ethylene terephthalate) (PET) resin, acrylonitrile-butadiene-styrene (ABS) resin and nylon resin, and method for the production thereof. In more detail, it relates to a binder resin composition for paint, printing ink, adhesive or primer that exhibits excellent adherence and other physical properties to sheets, films and moldings of these polyolefin, poly(vinyl chloride), polycarbonate, PET, ABS and nylon.

Background technologies

[0002] Because of many advantages of light weight, antirust, broad degree of freedom for design, etc., in recent years, plastics are used extensively as materials for automotive parts, electrical parts, building materials, etc. Above all, because of low price and many excellent properties such as moldability, chemical resistance, heat resistance, water resistance, good electrical characteristics, polyolefinic resins are used in a wide range as industrial materials and the growth of demand is also expected in future. Different from synthetic resins with polarity, however, polyolefinic resin is nonpolar and crystalline, hence it has also a drawback of difficult adhesion with it.

[0003] For this reason, as a pretreatment for painting and adhesion, method of providing plasma treatment or gas flame treatment onto the surface of moldings to activate, or method of painting primer (undercoating agent) having chlorinated polyolefin as a major component is adopted.

[0004] For the painting of polypropylene bumper of automobiles, for example, primer compositions having chlorinated modified polyolefin as a major component are disclosed in Japanese Unexamined Patent Publication No. Sho 57-36128 and Japanese Patent Publication No. Sho 63-36624.

[0005] With these primers comprising chlorinated products, the adherence onto polyolefin having been produced hitherto is excellent, but, it is the present situation that, with the advent of high-modularization of prime material and low-temperature baking in recent years, prime materials that give insufficient adherence are seen, making it impossible to correspond sufficiently with conventional chlorinated polyolefinic resins.

[0006] Furthermore, it is the present situation that the chlorinated polyolefinic resins exhibit relatively excellent adherence onto polypropylene prime material, but the adherence is insufficient onto other prime materials (poly(vinyl chloride), polycarbonate, PET, ABS and nylon).

[0007] The chlorinated polyolefin resins etc. having been proposed hitherto were those having chlorinated isotactic polypropylene (hereinafter IPP), wherein IPP produced by using Ziegler-Natta catalyst as a polymerization catalyst was chlorinated, as a major component.

[0008] Whereas, an adhesive that uses chlorinated syndiotactic polypropylene (hereinafter SPP), wherein SPP produced by using metallocene compound as a polymerization catalyst was chlorinated is disclosed (Japanese Patent No. 3045498 and Japanese Unexamined Patent Publication No. Hei 7-18016). However, this chlorinated SPP has more improved solvent solubility than that of conventional chlorinated IPP produced by using Ziegler-Natta catalyst as a polymerization catalyst, but it exhibited excellent adherence only when the prime material is polypropylene and it has a drawback of insufficient adherence onto other prime materials (poly(vinyl chloride), polycarbonate, PET, ABS and nylon).

[0009] The purpose of the invention is to provide a binder resin composition with good adherence and gasohol resistance to various polyolefins including polypropylene prime material, poly(vinyl chloride), polycarbonate, PET, ABS and nylon prime materials and also with excellent solvent solubility.

Disclosure of the invention

50 [0010] The inventors have come to solve the problems aforementioned with a binder resin composition characterized by being chlorinated propylenic random copolymer with weight average molecular weight of 3000 to 250000, wherein propylenic random copolymer with melting point (Tm) measured by differential scanning calorimeter (DSC) of 115 to 165°C obtained by copolymerizing propylene with other α-olefin in the coexistence of metallocene type catalyst was chlorinated to chlorine content of 10 to 40% by weight, after thermal degradation or directly without thermal degradation, and/or a binder resin composition characterized by being carboxyl group-containing chlorinated propylenic random copolymer with weight average molecular weight of 30000 to 220000, wherein propylenic random copolymer with melting point (Tm) measured by differential scanning calorimeter (DSC) of 115 to 165°C obtained by copolymerizing propylene with other α-olefin in the coexistence of metallocene type catalyst was grafted with α,β-unsaturated carbox-

ylic acid or its anhydride, after thermal degradation or directly without thermal degradation, and then chlorinated to chlorine content of 10 to 40% by weight.

[0011] The propylenic random copolymer being raw material of the invention is a copolymer obtained by copolymerizing propylene being major component with other α -olefin using metallocene catalyst as a polymerization catalyst. Commercial products such as Wintech (from Nippon Polychem Co., Ltd.) can also be used.

[0012] For the other α -olefin being comonomer, at least one kind can be selected from ethylene or a group consisting of olefins with number of carbon atoms of 4 or more. As the olefins with number of carbon atoms of 4 or more, 1-butene, 1-hexene, 4-methyl-1-penten 1-octene, etc. can be mentioned. By using metallocene catalyst, the range of copolymerizable comonomers can be widened over Ziegler-Natta catalyst.

[0013] For the metallocene catalyst to be used in the invention, publicly known one can be used. Concretely, such catalyst obtainable by combining components (A) and (B), and, if need be, additionally (C) described below is preferable.

[0014] Component (A): Metallocene complex being a compound of transition metal that belongs to groups 4 to 6 in the periodic table having at least one of conjugate 5-membered ring ligand.

[0015] Component (B): Cocatalyst capable of activating said metallocene complex (A) by reacting compound (B) with metallocene complex (A) (ion-exchangeable stratified silicate).

[0016] Component (C): Organic aluminum compound.

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[0017] The propylenic random copolymer of the invention can be produced by publicly known methods (Japanese Unexamined Patent Publication No. 2001-206914 etc.). For example, while supplying propylene, ethylene and hydrogen into the reactor and while continuously adding alkyl aluminum and metallocene catalyst, the production is performed.

[0018] The propylenic random copolymer of the invention is preferable to have melting point (hereinafter Tm) measured by differential scanning calorimeter (hereinafter DSC) of 115 to 165°C. If higher than 165°C, then the solvent solubility decreases. If lower than 115°C, then the adherence onto prime materials decreases. More preferable is low-melting point propylenic random copolymer with 115 to 135°C. Besides, in the measuring method of Tm by DSC in the invention, the evaluation was made with both peak temperature of melting and end temperature of melting at the time when sample (ca. 5mg) was taken and molten for 5 minutes at 200°C, and, after lowering the temperature to 40°C at a rate of 10°C/min to crystallize, the temperature was raised further to 200°C at a rate of 10°C/min to melt, using DSO measurement apparatus from Seiko Co.

[0019] For the propylenic random copolymer of the invention, it doesn't matter whether one thermally degraded at a temperature above melting point and below 350°C in the presence of radical generator or one without thermal degradation is used solely or by mixing, employing Banbury mixer, kneader, extruder or the like. The radical generator to be used for the reaction can be selected appropriately from publicly known ones, but, in particular, organic peroxide type compound is desirable.

[0020] As said organic peroxide type compounds, for example, di-t-butyl peroxide, dicumyl peroxide, t-butylcumyl peroxide, benzoyl peroxide, dilauryl peroxide, cumene hydroperoxide, t-butyl hydroperoxide, 1,1-bis(t-butylperoxy)-3,5,5-trimethylcyclohexane, 1,1-bis(t-butylperoxy)cyclohexane, cyclohexanone peroxide, t-butylperoxy benzoate, t-butylperoxy isobutyrate, t-butylperoxy-3,5,5-trimethyl hexanoate, t-butylperoxy-2-ethyl hexanoate, t-butylperoxyisopropyl carbonate, cumylperoxy octoate, etc. are mentioned.

[0021] The chlorinated propylenic random copolymer of the invention can be obtained by introducing chlorine to said propylenic random copolymer. The chlorination reaction is conducted by dissolving said propylenic random copolymer into chlorine-based solvent such as chloroform, and then by blowing-in gaseous chlorine while irradiating ultraviolet rays or in the presence of said organic peroxide.

[0022] The chlorine content is optimum to be 10 to 40% by weight, preferably 15 to 30% by weight. If lower than this range, then the adherence onto various prime materials becomes better, but the solubility into organic solvent decreases. Also, If higher than this range, then the adherence onto various prime materials decreases. Besides, the chlorine content is a value measured according to JIS-K7229.

[0023] The weight average molecular weight (hereinafter Mw) of the chlorinated propylenic random copolymer to be used in the invention is 3000 to 250000. If under 3000, the cohesion of resin is insufficient, and, if over 250000, then the handlings of ink and adhesive are lowered, which is unpreferable. Besides, the Mw in the invention is a value measured by means of gel permeation chromatography (hereinafter GPC, standard substance: polystyrene resin).

[0024] Moreover, the carboxyl group-containing chlorinated propylenic random copolymer of the invention can be obtained by introducing α,β -unsaturated carboxylic acid and chlorine to said propylenic random copolymer and the production thereof is possible by two methods mentioned below; namely, a method (first method) wherein, after directly graft polymerizing α,β -unsaturated carboxylic acid or its anhydride onto thermally degraded propylenic random copolymer or propylenic random copolymer without thermal degradation, the chlorination reaction is conducted, and a method (second method) wherein, after the chlorination reaction, α,β -unsaturated carboxylic acid or its anhydride is graft polymerized.

[0025] In following, concrete methods for the production will be exemplified. In the first method, first the method of directly graft copolymerizating α,β -unsaturated carboxylic acid or its anhydride onto thermally degraded propylenic random copolymer or propylenic random copolymer without thermal degradation can be performed by publicly known methods of a method (melt method) wherein said rein is molten by heating above melting point in the presence of radical generator to react, a method (solution method) wherein said rein is dissolved into organic solvent and then the solution is stirred under heating in the presence of radical generator to react, and the like.

[0026] In the case of the melt method, the reaction is conducted for a short time at a temperature above melting point and below 350°C employing Banbury mixer, kneader, extruder or the like, hence it has an advantage of simple manipulation.

[0027] On the other hand, in the case of the solution method, aromatic solvent such as toluene or xylene is used desirably as an organic solvent, but, besides, it may also be safe to use ester solvent, ketonic solvent or the like by mixing partially. The radical generator to be used for the reaction can be selected appropriately from publicly known ones, but, in particular, organic peroxide type compound is desirable and compounds listed above can be used.

[0028] However, in the case of the second method, when conducting the chlorination reaction after α,β -unsaturated carboxylic acid or its anhydride was graft copolymerized, it is required to replace said solvent with chlorinating solvent such as chloroform, hence, in the first method, melt method is more preferable.

[0029] Said chlorination reaction to be conducted after carboxyl group modification is performed by dissolving propylenic random copolymer graft copolymerized with α,β -unsaturated carboxylic acid or its anhydride into chlorine-based solvent such as chloroform and then by blowing-in gaseous chlorine while irradiating ultraviolet rays or in the presence of said organic peroxide.

[0030] In the method wherein, after the chlorination reaction, α,β -unsaturated carboxylic acid or its anhydride is graft polymerized being the second method, first, propylenic random copolymer is dissolved into chlorine-based solvent such as chloroform and the chlorination reaction is conducted similarly to the first method to produce chlorinated propylenic random copolymer, then solvent is changed to a solvent such as toluene or xylene, and α,β -unsaturated carboxylic acid or its anhydride is graft copolymerized in the presence of said organic peroxide. The reaction can be carried out at a temperature above 50°C and below the boiling point of solvent. However, in the second method, the graft polymerizability of α,β -unsaturated carboxylic acid or its anhydride is low at the reaction temperature above 50°C and below 100°C, and the chlorinated propylenic random copolymer has a potential to cause the de-hydrochloric acid above 100°C and below the boiling point of solvent. Hence the first method is more preferable than the second method. [0031] The purpose of graft copolymerizing α,β -unsaturated carboxylic acid or its anhydride onto propylenic random copolymer in the first method is to afford the adherence to upper coating paint when using the binder resin composition of the invention as a primer. The chlorinated polyolefin has originally low polarity, hence, when using as a primer (undercoating agent) as it is, the adherence onto PP prime material is good, but little adherence is seen onto high-polar upper coating paints (e.g. polyurethane paint and melamine paint). Hence it becomes important to enhance the polarity of chlorinated polyolefin by graft copolymerizing α,β -unsaturated carboxylic acid or its anhydride. As usable

polar upper coating paints (e.g. polyurethane paint and melamine paint). Hence it becomes important to enhance the polarity of chlorinated polyolefin by graft copolymerizing α,β -unsaturated carboxylic acid or its anhydride. As usable α,β -unsaturated carboxylic acids or their anhydrides, for example, maleic acid, citraconic acid, itaconic acid, aconitic acid and their anhydrides, acrylic acid, methacrylic acid, fumaric acid, mesaconic acid, etc. are exemplified, but, when considering the grafting ability onto polyolefin resin, maleic anhydride is most suitable. [0032] In the invention, the amount to be introduced by graft-copolymerizing α,β -unsaturated carboxylic acid or its

[0032] In the invention, the amount to be introduced by graft-copolymerizing α, β -unsaturated carboxylic acid or its anhydride is optimum to be 0 to 20% by weight and preferable is 0 to 10% by weight. If over 10% by weight, the moisture resistance tends to decrease when using as a primer.

[0033] The chlorine content is optimum to be 10 to 40% by weight, preferably 15 to 30% by weight. If lower than this range, then the adherence onto various prime materials becomes better, but the solubility into organic solvent decreases. Also, If higher than this range, then the adherence onto various prime materials decreases. Besides, the chlorine content is a value measured according to JIS-K7229.

[0034] The weight average molecular weight (hereinafter Mw) of the chlorinated propylenic random copolymer and the carboxyl group-containing chlorinated propylenic random copolymer to be used in the invention is 3000 to 250000 in the former. If under 3000, the cohesion of resin is insufficient, and, if over 250000, then the spraying property decreases, which is unpreferable. Also in the case of the latter, it is 30000 to 220000. If under 30000, the cohesion of resin is insufficient, and, if over 220000, then the spraying property decreases, which is unpreferable. Besides, the Mw in the invention is a value measured by means of gel permeation chromatography (hereinafter GPC, standard substance: polystyrene resin).

[0035] The chlorinated polyolefin accompanies the de-hydrochloric acid to degrade when exposing to ultraviolet rays or high temperature. When the chlorinated polyolefin causes the degradation by de-hydrochloric acid, decreased physical properties such as decreased adherence onto polypropylene prime material as well as coloring of resin, and aggravation of working environment due to freed hydrochloric acid are caused. From these facts, it is required to add a stabilizer. For obtaining this effect, it is preferable to add 0.1 to 5% by weight of stabilizer based on the resin component (solids). As the stabilizer, epoxy compound can be exemplified. The epoxy compound is not particularly restricted, but

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one compatible with chlorinated resin is preferable and such epoxy compound with epoxy equivalent of around 100 to 500 and with one or more epoxy groups in a molecule can be exemplified. For example, epoxidated soybean oil and epoxidated linseed oil epoxidated natural vegetable oils with unsaturated group with peracid such as peracetic acid, epoxidated fatty acid esters epoxidated unsaturated fatty acids such as oleic acid, tall oil fatty acid and soybean oil fatty acid, epoxidated alicyclic compounds represented by epoxidated tetrahydrophthalate, condensation products of bisphenol A and polyhydric alcohols with epichlorohydrin, for example, bisphenol A glycidyl ether, ethylene glycol glycidyl ether, propylene glycol glycidyl ether, glycerol polyglycidyl ether, sorbitol polyglycidyl ether, etc. can be exemplified. Moreover, monoepoxy compounds represented by butyl glycidyl ether, 2-ethylhexyl glycidyl ether, decyl glycidyl ether, stearyl glycidyl ether, allyl glycidyl ether, phenyl glycidyl ether, sec-butylphenyl glycidyl ether, tert-butylphenyl glycidyl ether, phenol polyethylene oxide glycidyl ether, etc. are exemplified. Moreover, metallic soaps such as calcium stearate and lead stearate used as stabilizers for poly(vinyl chloride) resin, organometallic compounds such as dibutyl tin dilaurate and dibutyl maleate and hydrotalcite compounds can also be used, and it may be safe to use these in combination.

[0036] The composition of the invention can also be used after dissolved into organic solvent. The solution concentration may be selected appropriately depending on the uses, but, the coating workability is injured at either too high or too low solution concentration, hence the resin concentration is preferable to be 5 to 60% by weight. For the solvent to be used, aromatic solvents such as toluene and xylene are preferable, and, besides, ester solvents such as ethyl acetate and butyl acetate, ketonic solvents such as methyl ethyl ketone and methyl isobutyl ketone, aliphatic solvents such as n-hexane and heptane, alicyclic solvents such as cyclohexane, methylcyclohexane and ethylcyclohexane can be used. In addition, for enhancing the preservative stability of resin solution, it is preferable to add alcohols such as methanol, ethanol and isopropyl alcohol, propylenic glycol ethers such as propylene glycol methyl ether, propylene glycol ethyl ether and propylene glycol tertiary-butyl ether solely or by mixing two or more kinds in amounts of 1 to 20% by weight based on said solvent. The recipe when dissolving the binder resin composition of the invention into organic solvent is possible by converting the chlorinating solvent such as chloroform being reaction solvent to said solvent using the difference of boiling point. Also, it may be safe that, after epoxy compound etc. were added to the reaction liquor having finished the reaction as stabilizers, it is fed to a vent extruder equipped with solvent-removing suction portion at screw shaft section to solidify and then dissolved into said solvent. The solidification can be carried out by the methods already known publicly, for example, by using vent extruder equipped with underwater cutting pelletizer at blow-off opening portion of extruder, vent extruder and pelletizer that cuts strand-like resin, etc.

[0037] The binder resin composition according to the invention can be used as a paint, printing ink, adhesive and primer applicable to films, sheets and moldings of polyolefin, poly(vinyl chloride), polycarbonate, PET, ABS and nylon. It may be used by coating as it is, but solvent, pigment and other additives may be added within a range not injuring the effect of the invention. Moreover, although said composition exhibits balanced physical properties of coated film by itself, it may be safe to use by further adding cyclized rubber, petroleum resin, cumarone-indene resin, chlorinated polyolefin resin, acrylic resin, alkyd resin, etc., if need be. In particular, it is preferable to contain the binder resin composition of the invention in amounts of 30% or more by weight.

Embodiment to put the invention into practice

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- 40 [0038] One of the features of the propylenic random copolymer produced by using metallocene catalyst as a polymerization catalyst is that it has a lower melting point glass transition point than that of the propylenic random copolymer produced by using conventional Ziegler Natta catalyst as a polymerization catalyst. It is considered therefore that, even if making the chlorine content low, the solvent solubility is good, and additionally, due to lower melting point, the adherence at the time of low-temperature baking becomes good.
- [0039] Moreover, it is mentioned that, as a feature of the propylenic random copolymer produced by using metal-locene catalyst as a polymerization catalyst, the molecular weight distribution is very narrow (Mw/Mn = ca. 2 or less).
 [0040] Furthermore, it has also newly become clear that, in the case of the propylenic random copolymer produced by using conventional Ziegler-Natta catalyst as a polymerization catalyst, decrease in the molecular weight, that is, formation of low-molecular weight components is caused inevitably, but, in the case of the propylenic random copolymer produced by using metallocene catalyst as a polymerization catalyst, formation thereof is little, because organic peroxide etc. are used when graft copolymerizing α,β-unsaturated carboxylic acid or its anhydride.
 - [0041] In the invention, although no distinct reason is seen, it has been found that, different from the propylenic random copolymer produced by using conventional Ziegler-Natta catalyst as a polymerization catalyst, the propylenic random copolymer produced by using metallocene catalyst as a polymerization catalyst exhibits excellent adherence onto wide range of prime materials such as poly(vinyl chloride), polycarbonate, PET, ABS and nylon, besides polyolefinic prime materials. Moreover, it is considered that, by using the propylenic random copolymer with less low-molecular weight components, produced by using metallocene catalyst as a polymerization catalyst, the gasohol resistance became good.

[0042] In following, the invention will be illustrated concretely based on examples, but the invention is not confined thereto.

[Example-1]

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[0043] A propylenic random copolymer (propylene ca. 97% - ethylene ca. 3%) (from Nippon Polychem Co., Ltd., MFR = 2.0g/10min, Tm = 125°C) produced by using metallocene catalyst as a polymerization catalyst was fed into a double shaft extruder set at a barrel temperature of 350°C to degrade thermally, thus obtaining a propylenic random copolymer with melt viscosity at 190°C of about 1500mPa-s. In a glass-lined reactor were put 500g of this resin. After 5L of chloroform were added, gaseous chlorine was blown-in from the bottom of reactor under a pressure of 2kg/cm², while irradiating ultraviolet rays to chlorinate. On the way, 3-point samplings were made and chloroform being solvent was distilled off by evaporator. Thereafter, replacement was made with toluene/cyclohexane = 70/30 (weight ratio) and Epicote 828 (from Yuka-Shell Epoxy Co., Ltd.) was added in amounts of 2% by weight based on resin as a stabilizer to obtain resin solutions of chlorinated propylenic random copolymer with resin concentration of 20% by weight. The physical properties of the resins are shown in Table 1. The resin solutions obtained were allowed to stand for one month at room temperature, but no changes were seen in the liquid state and appearance.

[Example-2]

[0044] In a glass-lined reactor were put 500g of propylenic random copolymer (propylene ca. 97% - ethylene ca. 3%) (Wintech, fromNippon Polychem Co., Ltd., MFR = 2.0g/10min, Tm = 125°C) produced by using metallocene catalyst as a polymerization catalyst. After 5L of chloroform were added, gaseous chlorine was blown-in from the bottom of reactor under a pressure of 2kg/cm², while irradiating ultraviolet rays to chlorinate. On the way, 3-point samplings were made and chloroform being solvent was distilled off by evaporator. Thereafter, replacement was made with toluene/cyclohexane = 70/30 (weight ratio) and Epiol TB (from Nippon Oil and Fats Co., Ltd.) was added in amounts of 2% by weight based on resin as a stabilizer to obtain resin solutions of chlorinated propylenic random copolymer with resin concentration of 20% by weight. The physical properties of the resins are shown in Table 1. The resin solutions obtained were allowed to stand for one month at room temperature, but no changes were seen in the liquid state and appearance.

[Example-3]

[0045] A propylenic random copolymer (propylene ca. 97% - ethylene ca. 3%) (from Nippon Polychem Co., Ltd., MFR = 2.0g/10min, Tm = 125°C) produced by using metallocene catalyst as a polymerization catalyst was fed into a double shaft extruder set at a barrel temperature of 350°C to degrade thermally, thus obtaining a propylenic random copolymer with melt viscosity at 190°C of about 2000mPa·s. In a four-neck flask attached with stirrer, condenser, thermometer and dropping funnel, 500g of this resin was molten by heating at 190°C. After nitrogen replacement in flask was performed for 10 minutes, 25g of maleic anhydride were put-in over about 5 minutes while stirring, and 2g of di-t-butyl peroxide were added dropwise over about 30 minutes as a radical generator. After the reaction was continued further for 30 minutes, unreacted maleic anhydride was removed while reducing the pressure in flask with aspirator. Next, this product was put in a glass-lined reactor, and, after 5L of chloroform were added, gaseous chlorine was blown-in from the bottom of reactor under a pressure of 2kg/cm², while irradiating ultraviolet rays to chlorinate. On the way, samplings were made and chloroform being solvent was distilled off by evaporator, respectively. Then, replacement was made with toluene/cyclohexane = 70/30 (weight ratio) to obtain 20% by weight solutions of chlorinated propylenic random copolymer modified with maleic anhydride. Epicote 828 (from Yuka-Shell Epoxy Co., Ltd.) was added in amounts of 4% by weight based on resin as a stabilizer. The physical properties of the resins are shown in Table 1. The resin solutions obtained were allowed to stand for one month at room temperature, but no changes were seen in the liquid state and appearance.

50 [Example-4]

[0046] In a double shaft extruder with L/D = 34 and ϕ = 40mm were put 500g of propylenic random copolymer (propylene ca. 97% - ethylene ca. 3%) (from Nippon Polychem Co. , Ltd., MFR = 7.0g/10min, Tm = 125°C) produced by using metallocene catalyst as a polymerization catalyst, 500g of maleic anhydride and 15g of dicumyl peroxide. The reaction was conducted making the retention time 10 minutes and barrel temperature 180°C (first barrel through seventh barrel) and deaeration was made at the seventh barrel to remove unreacted maleic anhydride, thus obtaining maleic anhydride-modified propylenic random copolymer. In a glass-lined reactor were put 500g of this resin, and, after 5L of chloroform were added, gaseous chlorine was blown-in from the bottom of reactor under a pressure of 2kg/cm²,

while irradiating ultraviolet rays to chlorinate. On the way, samplings were made and chloroform being solvent was distilled off by evaporator, respectively. Then, replacement was made with toluene/cyclohexane = 70/30 (weight ratio) to obtain 20% by weight solutions of chlorinated propylenic random copolymer modified with maleic anhydride. Epiol SB (from Nippon oil and Fats Co., Ltd.) was added respectively in amounts of 4% based on resin as a stabilizer. The physical properties of the resins are shown in Table 1. The resin solutions obtained were allowed to stand for one month at room temperature, but no changes were seen in the liquid state and appearance.

[Comparative example-1]

[0047] An isotactic polypropylene (IPP) produced by using Ziegler-Natta catalyst as a polymerization catalyst was fed into a double shaft extruder set at a barrel temperature of 350°C to degrade thermally, thus obtaining an IPP with melt viscosity at 190°C of about 2000mPa·s. Using 500g of this resin, resin solutions of chlorinated IPP with resin concentration of 20% by weight were obtained, similarly to Example 1. The physical properties of the resins are shown in Table 1. The resin solutions obtained were allowed to stand for one month at room temperature, but no changes were seen in the liquid state and appearance.

[Comparative example-2]

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[0048] Using 500g of syndiotactic polypropylene (SPP, MFR = 3. 7g/10min), resin solutions of chlorinated SPP with resin concentration of 20% by weight were obtained, similarly to Example 1. The physical properties of the resins are shown in Table 1. The resin solutions obtained were allowed to stand for one month at room temperature, but no changes were seen in the liquid state and appearance.

[Comparative example-3]

[0049] An isotactic polypropylene (IPP) produced by using Ziegler-Natta catalyst as a polymerization catalyst was fed into a double shaft extruder set at a barrel temperature of 350°C to degrade thermally, thus obtaining an IPP with melt viscosity at 190°C of about 2000mPa-s. Using 500g of this resin, resin solutions of maleic anhydride-modified chlorinated IPP with resin concentration of 20% by weight were obtained, similarly to Example 2. The physical properties of the resins are shown in Table 1. The resin solutions obtained were allowed to stand for one month at room temperature, but no changes were seen in the liquid state and appearance.

[Comparative example-4]

[0050] A syndiotactic polypropylene (SPP, MFR = 3.7g/10min) was fed into a double shaft extruder set at a barrel temperature of 350°C to degrade thermally, thus obtaining a SPP with melt viscosity at 190°C of about 2000mPa·s. From 500g of this resin, resin solutions of maleic anhydride-modified chlorinated SPP with resin concentration of 20% by weight were obtained, similarly to Example 2. The physical properties of the resins are shown in Table 1. The resin solutions obtained were allowed to stand for one month at room temperature, but no changes were seen in the liquid state and appearance.

[Comparative example-5]

[0051] In Example 1, without adding a stabilizer, resin solutions of chlorinated propylenic random copolymer with resin concentration of 20% by weight were obtained. When these resin solutions were allowed to stand for one month at room temperature, all of resin solutions discolored into reddish brown.

[Measuring method of physical properties of resin]

50 · MFR (Melt flow rate)

[0052] Measurement was made according to Melt Flow Rate in JIS-K-6758 Testing Method of Polypropylene (conditions: 230°C, load 2.16kgf).

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[0053] The evaluation was made with both peak temperature of melting and end temperature of melting at the time when sample (ca. 5mg) was taken and molten for 5 minutes at 200°C, and then, after lowering the temperature to 40°C

at a rate of 10°C/min to crystallize, the temperature was raised further to 200°C at a rate of 10°C/min to melt, using DSC measurement apparatus from Seiko Co.

· Chlorine content

[0054] Measurement was made according to JIS-K-7229.

- · Weight average molecular weight (Mw) and number average molecular weight (Mn)
- 10 [0055] Measurement was made by means of GPC (standard substance: polystyrene resin). Moreover, in the table, Mw/Mn shows the molecular weight distribution.

[Test of liquid state]

15 [0056] The resin solution obtained was allowed to stand for one month at room temperature, and the liquid state and appearance were evaluated visually.

Criterion for evaluation

20 [0057]

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- O: No changes in the liquid state and appearance, even after one month.
- Δ: Discoloration into reddish brown, after one month.
- ×: Gelation, after one month.

[Primer test]

[0058] After 100g of each of resin solutions (solids 20%) obtained from Examples 3 and 4 and Comparative examples 3 and 4 and 20g of titanium dioxide were kneaded for 3 hours in a sand mill, viscosity was adjusted with xylene so as to give 13 to 15 seconds/20°C through Ford cup No.4, and the mixture was painted onto an ultrahigh modulus PP plate (TX-933A, from Mitsubishi Chemical Industries Ltd.), poly(vinyl chloride) (PVC), polycarbonate (PC), PET, ABS or nylon-6 by air spray gun so as the film thickness to become about 10µm. Next, cure type two-component urethane paint was painted (film thickness ca. 30µm), which was dried for 30 minutes at 80°C and, after allowed to stand for 24 hours at room temperature, the physical properties were evaluated. The results of primer test are shown in Table 2.

· Adherence

[0059] On the coated surface, 100 cross-cuts that reach the base were made at intervals of 1 mm, and cellophane adhesive tape was adhered closely thereon. Then, it was peeled off in the direction of 180° to judge with the extent of remaining coated film.

- Gasohol resistance: The painted plate was soaked into regular gasoline/ethanol = 9/1 (v/v) for 120 minutes to
 observe the state of coated film.
 - Good: State of no abnormality in the coated film
 - No good: State of abnormality in the coated film
- Water resistance: The painted plate was soaked into warm water of 40°C for 240 hours to examine the state of coated film and adherence.
 - Good: Case of entirely no peeling
 - No good: Case of peeling caused

[Adhesiveness test]

- · Heat seal test
- [0060] Each of the resin solutions (solids 20 wt.%) obtained was coated onto an untreated PP, poly(vinyl chloride) (PVC) or PET with coating rod #14. After dried for 24 hours at room temperature, coated surfaces were superposed and heat sealed under the press conditions of 2 sec, 80°C and 1 kg/cm². After 24 hours, 180° peeling strength (g/cm) was measured with Tensilon (pulling speed: 50mm/min). The results of adhesiveness test are shown in Table 3.

[Ink test]

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[0061] After 100g of each of resin solutions (solids 20 wt.%) obtained and 20g of titanium dioxide were milled for 3 hours in a sand mill, the mixture was diluted with toluene so as the viscosity to become 25 to 30 seconds/20°C through #3 Zahn cup to prepare ink. With the ink obtained, adhesive tape peeling test and heat seal test were performed. The results of ink test are shown in Table 4.

· Adhesive tape peeling test

[0062] By the similar method to heat seal test, ink was coated onto an untreated PP, poly(vinyl chloride) (PVC) or PET. After dried for 24 hours at room temperature, cellophane adhesive tape was stuck onto ink-coated surface, and the state of coated surface when peeling off at a stroke was examined.

Criterion for evaluation

[0063]

Good: State of entirely no peeling No good: State of peeling caused

· Heat seal test

[0064] Same as adhesiveness test.

Table 1

	Ph	ysical properties of	resin and tes	t of liquid state		
	Chlorine content (wt.%)	Maleic anhydride (wt. %)(based on chlorinated resin)	Avera	Test of liquid state		
			Mw	Mn	Mw/Mn	
Ex.1-1	20.5		49800	26200	1.90	0
- 2	24.3	-	49700	26000	1.91	0
-3	32.0		49800	26200	1.90	0
Ex.2-1	19.5		190000	104000	1.83	0
- 2	25.0	-	191000	104000	1.83	0
-3	31.0		190000	103000	1.84	0
Ex.3-1	16.8	3.9	50800	27800	1.83	0
- 2	19.5	3.5	52400	27600	1.90	0
-3	26.1	3.3	53000	27400	1.93	0
Ex.4-1	16.0	5.0	120000	62500	1.92	0
- 2	20.8	5.3	125000	64000	1.95	0
-3	25.1	5.1	123000	62400	1.97	0
Comp.1-1	20.2		-	-	-	×
- 2	24.5	-	200000	68000	2.95	0
-3	31.5		210000	71000	2.94	0
Comp.2-1	20.2		198000	101500	1.95	0
- 2	30.5	-	198000	101000	1.96	0
-3	33.0		199000	101000	1.97	0
Comp.3-1	15.2	3.8	•	-	-	×
- 2	18.6	3.5	60000	27400	2.19	0

Table 1 (continued)

Physical properties of resin and test of liquid state								
	Chlorine content (wt.%)	Maleic anhydride (wt. %)(based on chlorinated resin)	Average molecular weight			Test of liquid state		
			Mw	Mn	Mw/Mn			
-3	26.0	3.4	60000	27400	2.19	0		
Comp.4-1	17.6	3.7	58000	29300	1.98	0		
- 2	22.6	3.4	58000	29100 '	1.99	0		
-3	26.5	3.3	57800	29200	1.98	0		
Comp.5-1	20.0		190000	104000	1.83	Δ		
- 2	25.5	-	191000	104000	1.83	Δ .		
- 3	31.5		190000	103000	1.84	Δ		

Note: With Comparative example 1-1 and Comparative example 3-1, the molecular weight could not be measured, since the resin solutions immediately after production were pudding-like.

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		_								_												_
5				Nylon	Adherence (/100)		100	100	100		100	100	100		1	0	0		20	20	0	dding-like.
10				ABS	Adherence (/100)		100	100	100		100	100	100			70	20		20	50	, 20	oduction was pu
15 20				PET	Adherence (/100)		100	100	100		100	100	100		•	0	0		0	0	0	ediately after pr
25			ial	5	Adherence (/100)		100	100	100		100	100	100		•	30	10		50	0	0	esin solution imn
30	Table 2		Prime material	PVC	Adherence (/100)		100	100	100		100	100	100		•	50	30		70	09	20	ed out, since the
35					Water resistance		Good	Good	Good		Good	Good	Good		ı	Good	Good		Good	Good	Good	nple 3-1, each test could not be carried out, since the resin solution immediately after production was pudding-like
40				ЬР	Gasohol resistance		Good	Good	Good		Good	Good	Good		ı	Good	No good		Good	Good	Good	nple 3-1, each tes
45						_																ive exar
50		Results of primer test			Adherence (/100)		100	100	100		100	100	100		1	100	100		100	100	100	Note: With Comparative exan
55		Results of				Ĕ.	3-1	-2	- 3	EX.	4-1	-2	- 3	Comp.	3-1	-,2	ę.	Сошр.	3-1	Ņ	ę-	Note:

Table 3

			Heat seal strength (g/cm)	
		PP prime material	PVC prime material	PET prime material
	Ex.1-1	860	830	830
1	- 2	780	770	750
	- 3	690	710	650
	Ex.2-1	900	820	780
	- 2	880	700	750
	- 3	730	680	660
	Ex.3-1	1200	1050	1100
	- 2	1000	1000	1050
	- 3	850	920	900
	Ex.4-1	1300	1350	1400
	- 2	1100	1200	1380
	- 3	900	1140	1220
	Comp.1-1	•	-	-
]	- 2	500	30	0
	- 3	400	0	0
	Comp.2-1	600	200	50
	- 2	500	100	30
	- 3	350	0	0
	Comp.3-1	-	-	-
	- 2	700	50	0
	- 3	600	0	0
	Comp.4-1	1000	30	10
	- 2	800	10	0
	- 3	650	0	0
	Comp.5-1	780	400	300
1	- 2	680	200	280
ĺ	- 3	600	180	100

Note: With Comparative example 1-1 and Comparative example 3-1, the tests could not be carried out, since the resin solutions immediately after production were pudding-like.

Table 4

	Peel	ing with adhesiv	e tape	Heat seal strength (g/cm)			
	PP prime material	PVC prime material	PET prime material	PP prime material	PVC prime material	PET prime material	
Ex.1-1	Good	Good	Good	650	680	630	
- 2	Good	Good	Good	590	600	530	
- 3	Good	Good	Good	450	460	420	
Ex.2-1	Good	Good	Good	700	720	750	
- 2	Good	Good	Good	500	700	700	
- 3	Good	Good	Good	400	660	650	

Table 4 (continued)

	Peel	ing with adhesiv	e tape	Heat seal strength (g/cm)			
	PP prime material	PVC prime material	PET prime material	PP prime material	PVC prime material	PET prime material	
Ex.3-1	Good	Good	Good	900	1000	1050	
- 2	Good	Good	Good	850	940	980	
- 3	Good	Good	Good	700	880	950	
Ex.4-1	Good	Good	Good	1000	1100	1200	
- 2	Good	Good	Good	920 ,	1050	1100	
- 3	Good	Good	Good	780	1000	1150	
Comp.1-1	-	-	-	-	-	-	
- 2	Good	No good	No good	400	0	0	
- 3	Good	No good	No good	320	0	0	
Comp.2-1	Good	No good	No good	580	100	30	
- 2	Good	No good	No good	410	40	0	
- 3	Good	No good	No good	290	0	0	
Comp.3-1	-	-	-	-	-	-	
- 2	Good	No good	No good	530	0	0	
- 3	Good	No good	No good	440	0	0	
Comp.4-1	Good	No good	No good	680	0	0	
- 2	Good	No good	No good	510	0	0	
- 3	Good	No good	No good	450	0	0	
Comp.5-1	Good	Good	Good	540	200	200	
- 2	Good	Good	Good	480	50	250	
- 3	No good	No good	No good	230	l o	0	

the resin solutions immediately after production were pudding-like.

Utilizability in the industry

[0065] From Table 1, the chlorinated products of propylenic random copolymer produced by using metallocene catalyst as a polymerization catalyst have good liquid state, even if the chlorine content may be low. Moreover, from Table 2, the chlorinated products of propylenic random copolymer produced by using metallocene catalyst as a polymerization catalyst have better gasohol resistance than that of the chlorinated products of IPP produced by using conventional Ziegler-Natta catalyst as a polymerization catalyst. Furthermore, from Tables 2 through 4, the chlorinated products of propylenic random copolymer produced by using metallocene catalyst as a polymerization catalyst exhibit good adherence not only onto polypropylene prime material, but also onto prime materials of poly(vinyl chloride), polycarbonate, PET, ABS, nylon, etc. It can be seen therefore that the binder resin compositions containing these chlorinated propylenic random copolymers and/or carboxyl group-containing chlorinated propylenic random copolymers are useful in the industry and, in particular, effective for paint, adhesive, heat sealing agent, printing ink and primer.

Claims

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- 1. A binder resin composition characterized by containing chlorinated propylenic random copolymer with weight average molecular weight of 3000 to 250000, wherein propylenic random copolymer obtained by copolymerizing propylene with other α-olefin in the coexistence of metallocene type catalyst is chlorinated to chlorine content of 10 to 40% by weight, stabilizer and organic solvent.
- 2. A binder resin composition, wherein the chlorinated propylenic random copolymer of Claim 1 is a carboxyl group-

containing chlorinated propylenic random copolymer with weight average molecular weight of 30000 to 220000, graft polymerized with α,β -unsaturated carboxylic acid or its anhydride in amounts of 0 to 20% by weight and then chlorinated to chlorine content of 10 to 40% by weight, or chlorinated to chlorine content of 10 to 40% by weight and then graft polymerized with α,β -unsaturated carboxylic acid or its anhydride in amounts of 0 to 20% by weight.

- 3. The binder resin composition of Claim 1 or 2, wherein the propylenic random copolymer has a melting point (Tm) measured by differential scanning calorimeter (DSC) of 115 to 165°C.
- 4. A method of producing binder resin composition of Claim 1 or 2, using the chlorinated propylenic random copolymer, wherein propylenic random copolymer with melting point (Tm) measured by differential scanning calorimeter (DSC) of 115 to 165°C obtained by copolymerizing propylene with other α-olefin in the coexistence of metallocene type catalyst is chlorinated to chlorine content of 10 to 40% by weight, after thermal degradation or without thermal degradation.
- 5. A method of producing binder resin composition of Claim 4, wherein the chlorinated propylenic random copolymer is a carboxyl group-containing chlorinated propylenic random copolymer graft polymerized with α,β-unsaturated carboxylic acid or its anhydride in amounts of 0 to 20% by weight and then chlorinated to chlorine content of 10 to 40% by weight, or chlorinated to chlorine content of 10 to 40% by weight and then graft polymerized with α,β-unsaturated carboxylic acid or its anhydride in amounts of 0 to 20% by weight.
 - 6. A paint applicable to films, sheets and moldings of polyolefin, poly(vinyl chloride), polycarbonate, PET, ABS and nylon, having the binder resin composition of any of Claims 1 to 3 as an effective component.
- 7. A printing ink applicable to films, sheets and moldings of polyolefin, poly (vinyl chloride), polycarbonate, PET, ABS and nylon, having the binder resin composition of any of Claims 1 to 3 as an effective component.
 - 8. An adhesive applicable to films, sheets and moldings of polyolefin, poly (vinyl chloride), polycarbonate, PET, ABS and nylon, having the binder resin composition of any of Claims 1 to 3 as an effective component.
- 30 9. A heat sealing agent applicable to films, sheets and moldings of polyolefin, poly(vinyl chloride), polycarbonate, PET. ABS and nylon, having the binder resin composition of any of Claims 1 to 3 as an effective component.
 - 10. A primer applicable to films, sheets and moldings of polyolefin, poly (vinyl chloride), polycarbonate, PET, ABS and nylon, having the binder resin composition of any of Claims 1 to 3 as an effective component.

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INTERNATIONAL SEARCH REPORT

International application No.
PCT/JP02/13345

A. CLAS Int.	SIFICATION OF SUBJECT MATTER .Cl ⁷ CO8L23/28, CO8F8/22, CO9D	123/28, C09J123/28	
According	to International Patent Classification (IPC) or to both	national classification and IPC	
B. FIELD	S SEARCHED		
	locumentation searched (classification system followed		
Int.	Cl' C08L23/28, C08F8/22, C09D	123/28, C090123/28	
		.* *	
Documenta	tion searched other than minimum documentation to the	ne extent that such documents are included	in the fields searched
Electronic d	lata base consulted during the international search (nar	ne of data base and, where practicable, sea	rch terms used)
C. DOCU	MENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where a		Relevant to claim No.
X	1	per Industries Co.,	1-10
	Ltd.),		
	24 April, 2001 (24.04.01), All references		
	(Family: none)		
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		00/42103 A1	
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Y	JP 11-302324 A (Daiso Co., I		1-10
	02 November, 1999 (02.11.99)	<i>'</i>	
1	All references (Family: none)		
	(ramily: none)	f	
			ţ
× Furth	er documents are listed in the continuation of Box C.	See patent family annex.	
• Special	categories of cited documents:	"I" later document published after the inter	mational filing date or
"A" docume	ent defining the general state of the art which is not	priority date and not in conflict with th	e application but cited to
	red to be of particular relevance document but published on or after the international filing	understand the principle or theory unde "X" document of particular relevance; the o	
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cited to	establish the publication date of another citation or other	"Y" document of particular relevance; the c	laimed invention cannot be
	reason (as specified) ent referring to an oral disclosure, use, exhibition or other	considered to involve an inventive step combined with one or more other such	
means		combination being obvious to a person	skilled in the art
	ent published prior to the international filing date but later e priority date claimed	"&" document member of the same patent f	amily
	ctual completion of the international search	Date of mailing of the international search	
	pril, 2003 (02.04.03)	15 April, 2003 (15.	04.03)
	ailing address of the ISA/ nese Patent Office	Authorized officer	
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Form PCT/ISA/210 (second sheet) (July 1998)

INTERNATIONAL SEARCH REPORT

International application No.
PCT/JP02/13345

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Y	JP 5-222320 A (Mitsui Petrochemical Industries,	1-10
1	Ltd.), 31 August, 1993 (31.08.93), Full description (Family: none)	1-10
Y	EP 148346 A2 (Mitsui Petrochemical Industries, Ltd.), 17 July, 1985 (17.07.85), All references 6 JP 60-99138 A 6 US 4755553 A 6 JP 88036624 B2 6 EP 148346 B1	1-10

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